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2 **Growth of nitrite-oxidizing *Nitrospira* and ammonia-oxidizing**
3 ***Nitrosomonas* in marine recirculating trickling biofilter reactors**

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31 **Originality-significance statement**

32 Aerobic nitrite oxidation (NO_2^- to NO_3^-) yields much less Gibbs free energy than aerobic
33 ammonia oxidation (NH_3 to NO_2^-), while dominance of nitrite-oxidizing bacteria (NOB) over
34 ammonia-oxidizing bacteria (AOB) has been found in marine recirculating trickling biofilter
35 reactors. Specific mechanism responsible for the formation of this puzzling microbial
36 community has not been explored in detail, and the present study shed light on this subject.
37 The present study shows novel ecological aspects of nitrite-oxidizing *Nitrospira* and
38 ammonia-oxidizing *Nitrosomonas* proliferated in trickling biofilter reactors.

39 **Summary (190 words)**

40 Aerobic ammonia and nitrite oxidation reactions are fundamental biogeochemical reactions
41 contributing to the global nitrogen cycle. Although aerobic nitrite oxidation yields 4.8-folds
42 less Gibbs free energy (ΔG_r) than aerobic ammonia oxidation in the NH_4^+ -feeding marine
43 recirculating trickling biofilter reactors operated in the present study, nitrite-oxidizing and not
44 ammonia-oxidizing *Nitrospira* (sublineage IV) outnumbered ammonia-oxidizing
45 *Nitrosomonas* (relative abundance; 53.8% and 7.59%, respectively). CO_2 assimilation
46 efficiencies during ammonia or nitrite oxidation were $0.077 \mu\text{mol}^{-14}\text{CO}_2/\mu\text{mol-NH}_3$ and 0.053
47 to $0.054 \mu\text{mol}^{-14}\text{CO}_2/\mu\text{mol-NO}_2^-$, respectively, and the difference between ammonia and
48 nitrite oxidation was much smaller than the difference of ΔG_r . Free-energy efficiency of
49 nitrite oxidation was higher than ammonia oxidation (31-32% and 13%, respectively), and
50 high CO_2 assimilation and free-energy efficiencies were a determinant for the dominance of
51 *Nitrospira* over *Nitrosomonas*. Washout of *Nitrospira* and *Nitrosomonas* from the trickling
52 biofilter reactors was also examined by quantitative PCR assay. Normalized copy numbers of
53 *Nitrosomonas amoA* was 1.5- to 1.7-folds greater than *Nitrospira nxrB* and 16S rRNA gene in
54 the reactor effluents. *Nitrosomonas* was more susceptible for washout than *Nitrospira* in the
55 trickling biofilter reactors, which was another determinant for the dominance of *Nitrospira* in
56 the trickling biofilter reactors.

57 **Introduction**

58 Nitrification is a key microbial process in the global nitrogen cycle and also for biological
59 nitrogen removal from wastewater. In the nitrification process, ammonia is aerobically
60 oxidized to nitrite by aerobic ammonia-oxidizing bacteria and archaea (AOB and AOA,
61 respectively), and the formed nitrite is subsequently oxidized to nitrate by aerobic nitrite-
62 oxidizing bacteria (NOB). Phylogenetically diverse NOB such as *Nitrospira* (phylum
63 *Nitrospirota*), *Nitrospina* (*Nitrospinota*), *Nitrobacter*, *Nitrotoga*, *Nitrococcus*
64 (*Proteobacteria*), *Nitrolancea* (*Chloroflexota*) have been identified by culture-dependent and
65 -independent techniques (Daims *et al.*, 2016). The genus *Nitrospira* consists of
66 phylogenetically diverse members (*i.e.*, at least six phylogenetic sublineages) (Lebedeva *et*
67 *al.*, 2011), and *Nitrospira* population has been found from wide range of man-made and
68 natural ecosystems (Daims *et al.*, 2016). Additionally, the members of the *Nitrospira*
69 sublineage II has a unique metabolic capability, complete ammonia oxidation (comammox),
70 and comammox *Nitrospira* oxidize ammonia to nitrate via nitrite in a single cell (Daims *et al.*,
71 2015; van Kessel *et al.*, 2015). Population of comammox *Nitrospira* has been found in
72 freshwater and groundwater ecosystems, but rarely found in marine environments (Xia *et al.*,
73 2018).

74 Thermodynamically, aerobic ammonia oxidation yields larger Gibbs free energy than
75 aerobic nitrite oxidation; therefore, numerical dominance of AOB and/or AOA over NOB is

76 expected where nitrification proceeds. Indeed, stoichiometric and thermodynamic calculation
77 of nitrification processes has enabled to approximate population size and growth yields of
78 AOA and NOB in ocean (Zakem *et al.*, 2018; Zhang *et al.*, 2020). On the other hand,
79 numerical dominance of *Nitrospira* over AOA/AOB has been described in the recirculating
80 trickling filter reactors operated for marine aquacultures (Foesel *et al.*, 2008; Keuter *et al.*,
81 2011; 2017; Brown *et al.*, 2013). Another example is our previous study (Oshiki *et al.*, 2020),
82 where *Nitrospira* outnumbered AOB and AOA in NH₄⁺-feeding marine recirculating trickling
83 filter reactors (down-hanging sponge, DHS, reactors); 55%, 10% and <0.1% of total biomass,
84 respectively. Numerical dominance of *Nitrospira* over AOB/AOA in the DHS reactors was
85 somewhat surprising because the ΔG_r of aerobic ammonia oxidation was 4.8-folds higher in
86 the NH₄⁺-feeding DHS reactor than that of aerobic nitrite oxidation (**Supplementary text 1**).
87 However, the mechanism(s) responsible for the dominance of *Nitrospira* over AOB and AOA
88 in marine trickling filter reactors has not been explored in detail.

89 Consequently, the present study aimed to examine how *Nitrospira* outnumbered the
90 population of AOB (*i.e.*, *Nitrosomonas*) in the DHS reactors. The two DHS reactors were
91 operated with feeding of the inorganic seawater media containing NH₄⁺ or NO₂⁻ (designated
92 as NH₄⁺- and NO₂⁻-feeding DHS reactors, respectively) at 20°C, and metagenomic analyses
93 using the biomass retained in the reactors were performed to examine metabolic potentials of
94 *Nitrospira* and *Nitrosomonas*. Because genomic data only suggested metabolic potential, a

95 series of batch incubations were performed to examine CO₂ assimilation efficiencies, free-
96 energy efficiencies, and H₂ oxidation activity. CO₂ assimilation efficiencies were determined
97 by examining ¹⁴CO₂ incorporation into the biomass, and this approach has been used for
98 determining the yields of carbon fixation by nitrifiers (Glover *et al.*, 1985; Tsai and Tuovinen
99 1986; Bayer *et al.*, 2022) and for the biomass yields of anaerobic ammonia oxidizing bacteria
100 (Ali *et al.*, 2015; Awata *et al.*, 2015). Apart from the above metagenomic and physiological
101 experiments, washout of *Nitrosomonas* and *Nitrospira* in the NH₄⁺-feeding DHS reactor was
102 examined by determining the copy number of AOB *amoA*, *Nitrospira nxrB*, and *Nitrospira*
103 16S rRNA gene in the reactor effluents by quantitative PCR (qPCR) assay.

104

105 **Results**

106 **Metagenomic analysis of NH_4^+ - and NO_2^- -enriched biomass**

107 NH_4^+ - and NO_2^- -feeding DHS reactors were operated continuously for more than 1 y without
108 disturbances, and *Nitrospira* proliferated as a dominant population in both the operated
109 reactors as examined by fluorescence *in-situ* hybridization (**Fig. 1**). Metagenomic analyses
110 using the biomass collected from the NH_4^+ - or NO_2^- -feeding DHS reactors (designated as the
111 NH_4^+ - or NO_2^- -enriched biomass, respectively) were performed, and the 35.4 and 52.8 M
112 reads of 200-bp paired-end reads corresponding to 14.2 and 21.1 Gb were obtained from the
113 NH_4^+ - and NO_2^- -enriched biomass, respectively. Those sequence reads were assembled into
114 30 bacterial bins, which contained the 6 *Nitrospira* (NPIRA01 to NPIRA06 bins) and 2
115 *Nitrosomonas* (NMNS01 and NMNS02 bins) bins (**Table 1**).

116 Relative abundances and phylogeny of the obtained 30 bins were shown in **Table 1**
117 and **Fig. S1**, respectively. The relative abundances of the *Nitrospira* (especially, the
118 NPIRA01, NPIRA02, NPIRA04 bins) and *Nitrosomonas* bins (the NMNS02 bin) were much
119 higher than the other bins, indicating that *Nitrospira* and *Nitrosomonas* were the predominant
120 bacteria in the NH_4^+ - and/or NO_2^- -enriched biomass. Especially, the sum of the relative
121 abundances of the NPIRA bins were 53.8% and 72.2% in the NH_4^+ - and NO_2^- -enriched
122 biomass, indicating *Nitrospira* was highly abundant in both the biomass. As for *Nitrosomonas*
123 bins, the sum of the relative abundances of the NMSN bins were 7.59% and 0.62% in the

124 NH_4^+ - and NO_2^- -enriched biomass (**Table 1**), indicating *Nitrospira* outnumbered
125 *Nitrosomonas* in both the DHS reactors.

126 **Phylogeny and metabolic potential of NPIRA bins**

127 The NPIRA bins were affiliated into the *Nitrospira* sublineage IV (Lebedeva *et al.*, 2011)
128 (**Fig. 2**). The average nucleotide identity (ANI) values among the NPIRA bins were 71–89%
129 (**Table S1**), indicating each NPIRA bins represented different *Nitrospira* species (Richter and
130 Rosselló-Móra, 2006). The NPIRA03 and NPIRA06 bins were affiliated into the *Nitrospira*
131 *marina* clade including *Nitrospira marina*, a mesophilic and halophilic nitrite oxidizing
132 bacterium. Other NPIRA bins were affiliated into a phylogenetically-different clade in which
133 closely-relating *Nitrospira* genome was not available in public database (accessed on Jan.
134 2021). Phylogenetic affiliation of this clade was examined using the 16S rRNA gene sequence
135 located in the NPIRA02 bin (the NPIRA02_r00020 gene). The NPIRA02_r00020 gene
136 showed the 97.5% identity with the partial 16S rRNA gene sequences of *Candidatus*
137 *Nitrospira salsa* clone Cb18 (accession number KC706459.1) (**Fig. S2**), and this clade was
138 tentatively designated as the *Nitrospira salsa* clade in the present study. The NPIRA01,
139 NPIRA02, and NPIRA04 bins affiliated into the *Nitrospira salsa* clade were dominant
140 *Nitrospira* (*i.e.*, >10% of relative abundance in a biomass) both in the NH_4^+ - or NO_2^- -feeding
141 DHS reactors (**Table 1**).

142 Metabolic potential of the NPIRA bins was investigated by examining presence and
143 absence of functional genes (**Table S2**). The genes required for nitrite oxidation
144 (nitrite:nitrate oxidoreductase, *nxr*), energy conservation (cytochrome *bd*-like heme-copper
145 terminal oxidase), NAD(P)H generation (complex I), and CO₂ fixation through the rTCA
146 cycle (2-oxoglutarate:ferredoxin oxidoreductase, OGOR, five- or four-subunit
147 pyruvate:ferredoxin oxidoreductase, POR, and ATP-citrate lyase, ACL) were conserved in the
148 NPIRA bins (a full description is available as **supplemental text 2**). On the other hand,
149 orthologues of commamox *Nitrospira amoCAB* (threshold *e*-value of blastp search 10⁻¹⁵) were
150 not found in the NPIRA bins and also in the other known *Nitrospira* sublineage IV genomes.
151 The orthologue of commamox *Nitrospira hao* was also missing in the NPIRA bins and the
152 *Nitrospira* sublineage IV genomes, whereas the orthologue encoding octaheme cytochrome *c*
153 (a phylogenetically-relevant protein with Hao) (Bergmann *et al.*, 2005) found in the canonical
154 NO₂⁻-oxidizing *Nitrospira* genomes (*Nitrospira moscoviensis* and *Nitrospira japonica*
155 genomes) was located in the NPIRA2 and NPIRA3 bins (**supplemental text 2** and **Table S3**).

156 *Nitrospira* bacteria are metabolically versatile bacteria, and hydrogenotrophic growth
157 of *Nitrospira moscoviensis* (Koch *et al.*; 2014) has been demonstrated. The genes required for
158 H₂ oxidation and also degradations of carbohydrate and protein were conserved in the NPIRA
159 bins. A *hyd* gene cluster encoding a Group 3b NiFe hydrogenase and accessory proteins
160 required for the maturation of the NiFe hydrogenase was conserved in the NPIRA03 and

161 NPIRA04 bins (**Table S2**). On the other hand, the genes encoding putative Group 2a NiFe
162 hydrogenase (HupSL) were located in the NPIRA01 and NPIRA04 bins; therefore, the
163 NPIRA04 bin had both the Group 3b and Group 2a NiFe hydrogenase as previously found in
164 *Ca. Nitrospira alkalitolerans* (Daebeler *et al.*, 2020). The NPIRA03 and NPIRA06 bins had
165 the gene encoding formate dehydrogenase (Fdh) involved in formate oxidation. *Nitrospira*
166 *marina* cells can grow chemoorganotrophically on formate even without nitrite (Bayer *et al.*,
167 2021), and the NPIRA03_20570 and NPIRA06_25420 proteins showed 91.44 and 91.59%
168 identities with *Nitrospira marina* Fdh (the NMARINA_v1_1399 protein). On the other hand,
169 the *fdh* was not found in the NPIRA bins affiliated to *Nitrospira salsa* clade. The genes
170 encoding the enzymes involved in the degradation of carbohydrate (glycoside hydrolase, β -
171 glucosidase A, and alpha-amylase) and protein (secreted peptidases) were conserved in the
172 NPIRA03, NPIRA04, and NPIRA06 bins. As for the uptake of carbohydrate, amino acid and
173 peptides, ABC transporters for amino acid, oligo- and dipeptides were found in the NPIRA
174 bins whereas the sugar transporters were only found in specific NPIRA bins. The genes
175 encoding sugar ABC transporter were located in the NPIRA04 bin, whereas putative
176 carbohydrate-selective porin was found from NPIRA01, NPIRA02 and NPIRA03 bins.

177 **Phylogeny and metabolic potential of NMNS bins**

178 The NMNS01 and NMNS02 bins were affiliated into the *Nitrosomonas* sp. Nm143 and
179 *Nitrosomonas aestuarii/marina* clades, respectively (**Fig. 3**), and the members of

180 *Nitrosomonas* sp. Nm143 and *Nitrosomonas aestuarii* were previously found in recirculating
181 marine aquaculture systems (Itoi *et al.*, 2006; Foesel *et al.*, 2008). The genes required for
182 aerobic ammonia oxidation and energy conservation (*amo*, *hao*, complex III and terminal
183 oxidase) and for NAD(P)H generation (*i.e.*, reverse electron transport) were generally
184 conserved in the NMNS01 and NMNS02 bins (See **Supplemental text 2** for details.).
185 Orthologue of *amo* was not found from the obtained bins other than NMNS01 and NMNS02
186 bins. The NMNS bins had the genes encoding ribulose-1,5-bisphosphate carboxylase
187 (RuBisCO) and ribulose-5-phosphate kinase (**Table S4**), suggesting that those *Nitrosomonas*
188 fixed CO₂ using the Calvin-Benson cycle. It was previously shown that *Nitrosomonas* was
189 capable of H₂ oxidation coupled with nitrite reduction (Bock *et al.*, 1995), and the NMNS
190 bins had the genes encoding putative NiFe hydrogenase and accessory proteins (*i.e.*, Group 3b
191 and Group 3d NiFe hydrogenase for the NMNS01 and NMNS02 bins, respectively) (**Table**
192 **S4**). Those genes were not located as a single gene cluster but found as multiple gene clusters
193 as previously found in the *Nitrosomonas oligotropha* genome (Sedlacek *et al.*, 2019).

194 **Cultivation-dependent characterization of NH₄⁺- and NO₂⁻-enriched biomass**

195 ***CO₂ assimilation and free-energy efficiencies during aerobic ammonia or nitrite oxidation***

196 ¹⁴C₂ assimilation into the NH₄⁺- and NO₂⁻-enriched biomass during aerobic ammonia and
197 nitrite oxidation was examined by incubating the biomass with the addition of 0.5 mM
198 NH₄⁺ or NO₂⁻ and ¹⁴C-labeled sodium bicarbonate. The NH₄⁺-enriched biomass consumed

199 both NH_4^+ and NO_2^- , and produced NO_3^- stoichiometrically (**Fig. 4, left**). During the ammonia
200 and nitrite oxidation, $^{14}\text{CO}_2$ was assimilated into the NH_4^+ -enriched biomass, and the CO_2
201 assimilation efficiencies were determined to be 0.13 ± 0.019 (mean \pm standard deviation
202 derived from triplicate incubation) $\mu\text{mol-CO}_2/\mu\text{mol-NH}_3$ and $0.053 \pm 0.013 \mu\text{mol-CO}_2/\mu\text{mol-}$
203 NO_2^- , respectively (**Table 2**). It is notable that autotrophic bacteria release a part of fixed CO_2
204 as dissolved organic carbon (DOC) (Oshiki *et al.*, 2011), and the determined $^{14}\text{CO}_2$
205 assimilation does not include the fraction of DOC; *e.g.*, approx. 6-8% and 12% of fixed CO_2
206 were released as DOC in the culture of marine *Nitrosomonas* and *Nitrospira marina* Nb-295,
207 respectively (Bayer *et al.*, 2022). No $^{14}\text{CO}_2$ assimilation was detected from the pasteurized
208 biomass and the biomass incubated without the addition of NH_4^+ or NO_2^- . The CO_2
209 assimilation efficiency of the ammonia oxidation reaction could not be determined directly
210 because the formed NO_2^- was subsequently oxidized to NO_3^- without the accumulation of
211 NO_2^- (**Fig. 4, left**). Therefore, the CO_2 assimilation efficiency of the ammonia oxidation was
212 approximated by subtracting the CO_2 assimilation efficiency of ammonia oxidation reaction
213 with that of nitrite oxidation reaction (*i.e.*, $0.13 \mu\text{mol-CO}_2/\mu\text{mol-NH}_3$ and $0.053 \mu\text{mol-}$
214 $\text{CO}_2/\mu\text{mol-NO}_2^-$, respectively), which was $0.077 \mu\text{mol-CO}_2/\mu\text{mol-NH}_3$.

215 As for the NO_2^- -enriched biomass, the biomass consumed NO_2^- and
216 stoichiometrically produced NO_3^- (**Fig. 4, right**). On the other hand, the NO_2^- -enriched
217 biomass did not consume NH_4^+ , and no $^{14}\text{CO}_2$ assimilation was found. The CO_2 assimilation

218 efficiency of the nitrite oxidation reaction was determined to be $0.054 \pm 0.019 \mu\text{mol-}$
219 $\text{CO}_2/\mu\text{mol-NO}_2^-$, which was the same with that determined using the NH_4^+ -enriched biomass
220 (*i.e.*, $0.053 \pm 0.013 \mu\text{mol-CO}_2/\mu\text{mol-NO}_2^-$).

221 Free-energy efficiencies during the ammonia and nitrite oxidation were calculated
222 from the obtained CO_2 assimilation efficiencies (see the section Experimental procedures for
223 the calculation of the efficiencies), and compared with those previously determined using
224 AOB and NOB cultures (**Table 2**). The CO_2 assimilation and free-energy efficiencies of the
225 ammonia oxidation reaction obtained in the present study (*i.e.*, $0.077 \mu\text{mol-CO}_2/\mu\text{mol-NH}_3$
226 and 13%, respectively) were comparable with those previously determined from AOB
227 cultures. As for the CO_2 assimilation and free-energy efficiencies of the nitrite oxidation
228 reaction, those obtained in the present study (*i.e.*, 0.053 to $0.054 \mu\text{mol-CO}_2/\mu\text{mol-NO}_2^-$ and
229 31 to 32%, respectively) were greater than those previously determined from NOB including
230 *Nitrospira marina* Nb-295 (Bayer *et al.*, 2022). The 31 - 32% of the free-energy efficiencies
231 of the nitrite oxidation was >2.3-fold higher than that of ammonia oxidation reaction (*i.e.*,
232 13%).

233 *Activities of H₂ oxidation*

234 The above metagenomic analysis suggested that *Nitrospira* sp. NPIRA03, NPIRA01 and
235 NPIRA04 and *Nitrosomonas* sp. NSMS01 and NSMS02 were capable of H_2 oxidation, and
236 the activities of the H_2 oxidation of NH_4^+ - and NO_2^- -enriched biomass were examined by

237 performing batch incubations. It should be noted that the genes encoding putative NiFe
238 hydrogenase were also found from the bins other than NPIRA and NSMS bins (*i.e.*,
239 DHS20C07, DHS20C08, DHS20C16, DHS20C18, and DHS20C20 bins). Involvement of
240 those bacteria to H₂ oxidation could not be ruled out here, while the abundance of those bins
241 was much less than *Nitrospira* bins (**Table 1**). Although both the NH₄⁺- and NO₂⁻-enriched
242 biomass showed the activities of H₂ oxidation, the activities appeared after 4 d of incubation
243 (**Fig. 5**). Occurrence of the lag phase indicated that H₂ oxidation was not an active metabolic
244 pathway in the NH₄⁺- and NO₂⁻-enriched biomass at least in the operated DHS reactors. This
245 conclusion agreed with the observation obtained from the batch incubation in which the
246 NH₄⁺- and NO₂⁻-enriched biomass were incubated with the addition of ¹⁴CO₂ and H₂. The
247 NH₄⁺- and NO₂⁻-enriched biomass were incubated for 18 h (*i.e.*, within the above lag phase),
248 and no ¹⁴CO₂ assimilation was found in both the biomass during the 18 h of incubation.

249 *Washout of Nitrospira and Nitrosomonas from the NH₄⁺-feeding DHS reactor*

250 Washout of *Nitrospira* and *Nitrosomonas* cells from the NH₄⁺-feeding DHS reactor
251 were examined by determining the copy number of AOB *amoA*, *Nitrospira nxrB* and
252 *Nitrospira* 16S rRNA gene in reactor effluents. For the purpose, the qPCR assays of AOB
253 *amoA*, *Nitrospira nxrB* and *Nitrospira* 16S rRNA gene were carried out. There was a linear
254 relationship between the log copy number of standard DNAs and threshold cycle values (C_t
255 value) ($R^2 > 0.995$), and efficiencies of PCR amplification were 1.83 to 2.02. As shown in **Fig.**

256 6, all the AOB *amoA*, *Nitrospira nxrB* and *Nitrospira* 16S rRNA gene were detected from the
257 effluents, indicating both *Nitrosomonas* and *Nitrospira* were detached and washed out from
258 the NH₄⁺-feeding DHS reactor. The ratio of the copy numbers of AOB *amoA* to *Nitrospira*
259 *nxrB* and *Nitrospira* 16S rRNA gene increased in the effluents. Especially, *Nitrospira* 16S
260 rRNA gene was less abundant than AOB *amoA* in the effluents, indicating larger amounts of
261 *Nitrosomonas* population was washed out from the NH₄⁺-feeding DHS reactor. _____

262 **Discussion**

263 *Nitrospira* bacteria affiliated into the *Nitrospira salsa* or *Nitrospira marina* clade were
264 enriched in the NH_4^+ - and NO_2^- -feeding DHS reactors, and those *Nitrospira* outnumbered the
265 population of *Nitrosomonas* both in the DHS reactors. Those *Nitrospira* enriched in the DHS
266 reactors were most likely canonical nitrite-oxidizing *Nitrospira* because 1) the NPIRA bins
267 and the known *Nitrospira* sublineage IV genomes did not have the orthologue of *amoCAB*
268 and *hao* and 2) the *Nitrospira* of the NO_2^- -enriched biomass did not show the activity of
269 aerobic ammonia oxidation (**Fig. 4**) although they (*i.e.*, the NPIRA01 to NPIRA06) were
270 commonly found in the NH_4^+ - and NO_2^- -enriched biomass. Additionally, commamox
271 *Nitrospira* (affiliated into the *Nitrospira* sublineage II) has been found from freshwater and
272 groundwater ecosystems (Xia *et al.*, 2018), whereas no commamox *Nitrospira* has been
273 recognized from the *Nitrospira* sublineage IV often found in saline environments (Daims *et*
274 *al.*, 2016; Park *et al.*, 2020). It is obvious to raise the question of how *Nitrospira* sp.
275 NPIRA02 and NPIRA04 outnumbered *Nitrosomonas* sp. NMNS01 and NMNS02 in the
276 NH_4^+ -feeding DHS reactor. Not only the present study, the previous studies have also
277 indicated *Nitrospira* sublineage IV population outnumbered the population of aerobic

278 ammonia oxidizers in marine aquaculture systems (**Table S5**) (Foesel *et al.*, 2008; Keuter *et*
279 *al.*, 2011; 2017; Brown *et al.*, 2013).

280 Aerobic ammonia oxidation reaction yields 4.8-times higher free energy than NO_2^-
281 oxidation reaction in the NH_4^+ -feeding DHS reactor. However, the free energy recovered
282 from the ammonia oxidation reaction must be much lower than the ΔG_r due to the following
283 reasons. First, there is no evidence that the first reaction of aerobic ammonia oxidation to
284 NH_2OH involves translocation of H^+ and formation of proton motive force (Costa *et al.*, 2006;
285 Simon and Klotz, 2013). The reaction of aerobic ammonia oxidation to NH_2OH is a
286 monooxygenation reaction catalyzed by Amo (Lancaster *et al.*, 2018), and the free energy
287 released during monooxygenation reactions (more specifically, O_2 reduction reaction) are not
288 conserved and dissipated (VanBriesen, 2001). Dissipation of the free energy during the
289 monooxygenation reaction of CH_4 (Yuan and VanBriesen, 2002) and NH_3 (Hollocher *et al.*,
290 1982) has been described, and both the reactions are catalyzed phylogenetically relevant
291 monooxygenase (i.e., Pmo and Amo, respectively). Indeed, *Nitrosomonas europaea* cells
292 showed nearly same effective H^+/O ratios during aerobic ammonia and NH_2OH oxidation
293 (i.e., 4.1 and 3.9 of effective H^+/O ratios, respectively) (Hollocher *et al.*, 1982), indicating the
294 contribution of the reaction of ammonia oxidation to NH_2OH to the translocation of H^+ is
295 minor. The $\Delta G_r'^\circ$ for the reaction of aerobic ammonia oxidation to NH_2OH was -170.5 kJ
296 mol-NH_3^{-1} (**Supplementary text 1**), and this free energy (accounting for more than half of

297 ΔG_r 'o of aerobic ammonia oxidation) will be dissipated. Secondly, the oxidation of 1 mol
298 NH_2OH to NO_2^- releases 4 e^- , whereas the amounts of the e^- available for the respiration are
299 less than 2 e^- due to the following reasons; 1) the 2 e^- out of the produced 4 e^- is consumed to
300 oxidize 1 mol NH_3 to NH_2OH by Amo (Whittaker *et al.*, 2000), and 2) a part of the produced
301 4 e^- is consumed to generate NAD(P)H by reverse electron transport. In *Nitrosomonas*
302 *europaea* cells, the 0.35 e^- enters reverse electron flow (Whittaker *et al.*, 2000). Additionally,
303 the biochemistry of NH_2OH oxidation to NO_2^- by AOB is still controversial because the
304 *Nitrosomonas europaea* Hao oxidized NH_2OH to NO but not further to NO_2^- , and specific
305 mechanisms of NO oxidation to NO_2^- has not been elucidated (Carantoa and Lancaster, 2017;
306 Lancaster *et al.*, 2018). NO oxidation to NO_2^- releases 1 e^- , and the involvement of the
307 released e^- in the respiration of AOB needs to be investigated in other studies. The above
308 discussion pointed out that the amounts of the free energy recovered from aerobic ammonia
309 oxidation were much lower than that calculated as ΔG_r of the reaction. The reduction of the
310 free energy recovered from aerobic ammonia oxidation would result in the low free-energy
311 efficiencies of the ammonia oxidation reaction shown in **Table 2** (*i.e.*, 7 to 13%).

312 The ΔG_r of aerobic nitrite oxidation was much smaller than that of aerobic ammonia
313 oxidation ($-83.5 \text{ kJ mol-NO}_2^{-1}$ and $-283.3 \text{ kJ mol-NH}_3^{-1}$ at the batch incubation,
314 respectively,) (**Supplementary text 1**), while the CO_2 assimilation efficiency during aerobic
315 nitrite oxidation were close to that of the ammonia oxidation reaction; *i.e.*, $0.053 \mu\text{mol-}$

316 CO₂/μmol-NO₂⁻ (**Table 2**). Such high CO₂ assimilation and free-energy efficiencies (31-32%)
317 were rarely observed from axenic cultures of NOB (*e.g.*, *Nitrospira marina* Nb-295; 0.032
318 μmol-CO₂/μmol-NO₂⁻ and 22%, respectively), but high biomass yield of marine NOB
319 (*Nitrospinae*) was previously found in environmental samples (Kitzinger *et al.*, 2020) where
320 the NOB coexisted with other microbes. Low free-energy efficiencies of aerobic nitrite
321 oxidation found in the axenic cultures were reasonable because NO₂⁻ oxidation reaction can
322 not couple with the reduction of quinone molecules (ubiquinone_{ox/red}, ΔE° = +0.11 V) directly
323 due to the high redox potential of NO₂⁻/NO₃⁻ oxidation reaction (ΔE° = +0.42 V) (Madigan *et*
324 *al.*, 2019). Additionally, the electrons released from the NO₂⁻ oxidation reaction enter to
325 terminal oxidase bypassing a cytochrome *bc₁* complex (Lücker *et al.*, 2010; Simon and Klotz
326 *et al.*, 2013), resulting in the decrease of the number of translocated H⁺ during the respiration:
327 therefore, the free-energy efficiencies of nitrite oxidation reaction were expected to be low.
328 On the other hand, *Nitrospira* in the NH₄⁺- and NO₂⁻-feeding DHS reactors showed high CO₂
329 assimilation and free energy efficiencies. As compared with *Nitrobacter* and *Nitrococcus*, the
330 following bioenergetic advantages of the nitrite oxidation by *Nitrospira* were often introduced
331 in literatures; 1) nitrite oxidation occurs in the periplasmic spaces (Spieck *et al.*, 1998; Lücker
332 *et al.*, 2010), which directly contributes to the generation of proton motive force across a
333 membrane (*i.e.*, 1 NO₂⁻ + 1 H₂O → 1 NO₃⁻ + 2 H⁺_{periplasm} + 2 e⁻), and 2) *Nitrospira* used an
334 energetically more efficient rTCA cycle for CO₂ fixation as compared with *Nitrobacter* and

335 *Nitrosococcus* who used the Calvin-Benson cycle. However, the difference of CO₂ fixation
336 pathway might not result in a drastic change of CO₂ assimilation efficiency although the
337 Calvin-Benson cycle requires more ATPs investment for CO₂ fixation and involves a wasteful
338 oxygenase side reaction of ribulose-1,5-bisphosphate carboxylase/oxygenase (Berg 2011;
339 Bayer *et al.*, 2022). For the fixation of 3 mol CO₂ (*i.e.*, HCO₃⁻) to 1 mol of
340 phosphoglyceraldehyde which is the simplest sugar and a precursor for the biomass synthesis
341 (White *et al.*, 2012), the rTCA cycle requires 5 and 6 mol of ATP and NADPH equivalents
342 (such as NAD(P)H, ferredoxin and FADH₂), whereas the Calvin-Benson cycle requires 9 and
343 6 mol of ATP and NADPH equivalents (Bar-Even *et al.*, 2010). Therefore, the amounts of
344 NADPH equivalents are the same between the two pathways. To generate 6 mol of NAD(P)H
345 from NAD(P)⁺ by the reverse electron transport, *Nitrosomonas europaea* and *Nitrobacter*
346 *winogradskyi* consumed 30 mol of ATP (Aleem, 1966; Sewell and Aleem, 1969); therefore,
347 the energy cost for reverse electron flow is >3 folds higher than that for CO₂ fixation. The
348 ATP cost of the rTCA cycle was 4 mol-ATP/mol-phosphoglyceraldehyde fewer than that of
349 the Calvin-Benson cycle, while this energy conservation is much smaller than the energy
350 consumption for reverse electron transport. Indeed, the CO₂ assimilation efficiencies
351 previously determined from marine *Nitrococcus* with Calvin-Benson cycle (*Nitrococcus*
352 *mobilis*) were comparable with that of marine *Nitrospira* with rTCA cycle (*Nitrospira marina*

353 Nb-295); *i.e.*, 0.014-0.031 and 0.032 $\mu\text{mol-CO}_2/\mu\text{mol-NO}_2^-$, respectively (Glover 1985; Bayer
354 *et al.*, 2022) (**Table 2**).

355 It is notable that NOB can receive some micronutrients produced in microbial
356 community (Mee *et al.*, 2014; Kim *et al.*, 2021), which would raise CO₂ assimilation
357 efficiencies of *Nitrospira*. One example is the vitamin-B₁₂ auxotrophy of *Nitrospira marina*.
358 The *Nitrospira marina* genome lacked a couple of genes required for the biosynthesis of
359 vitamin B₁₂, and their growth ceased in vitamin-B₁₂ deficient media (Bayer *et al.*, 2021).
360 However, the growth of *Nitrospira marina* was found in a mixed culture even when vitamin
361 B₁₂ was not supplied into the media (Park *et al.*, 2020). Additionally, *Nitrospira* cells can
362 incorporate organic matters available in the culture. The addition of undefined organic matters
363 such as tryptone increased apparent growth yields of *Nitrospira marina* (Watson *et al.*, 1986;
364 Bayer *et al.*, 2021), and formate utilization by *Nitrospira* bacteria has been also demonstrated
365 (Gruber-Dorninger *et al.*, 2015; Koch *et al.*, 2015; Lawson *et al.*, 2021). Availability of
366 organic matters in the operated DHS reactors fed with inorganic media was suggested from
367 the growth of heterotrophic bacteria in the reactors. In the NH₄⁺- and NO₂⁻-feeding DHS
368 reactors, heterotrophs (*i.e.*, the DHS20C01 to DHS20C21 bins) accounted for *ca.* 10% of total
369 biomass (**Table 1**), and the presence of those heterotrophs indirectly indicated that organic
370 matters were available in the operated DHS reactors (likely in the form of extracellular
371 polymeric substances, soluble microbial products and cell debris). The obtained *Nitrospira*

372 bins (and also NMNS bins, **Table S6**) had a couple of (di)peptide transporters and ABC
373 transporters, and scavenging micronutrients might contribute to the increase of apparent CO₂
374 assimilation efficiencies. In addition to the possible interactions of micronutrients, oxidative
375 stress in mixed communities would be less because coexisting microbes can scavenge O₂ and
376 reactive oxygen species. Less oxidative stress would reduce energy demands of the rTCA
377 cycle of *Nitrospira* where oxygen-sensitive enzymes (*e.g.*, four-subunit pyruvate:ferredoxin
378 oxidoreductase) are involved (Berg *et al.*, 2011; Bayer *et al.*, 2022).

379 Although the above discussion provided a bioenergetic insight in the CO₂
380 assimilation of AOB and NOB, the growth yields of AOB were still higher than those of NOB
381 including *Nitrospira* (**Table 2**). How did *Nitrospira* with lower CO₂ assimilation efficiencies
382 outnumbered *Nitrosomonas* in the NH₄⁺-feeding DHS reactor? Notably, the DHS reactor and
383 the recirculating marine aquaculture systems in the previous studies were operated as trickling
384 filter reactors, and *Nitrospira* and AOB proliferated in the form of biofilm on the biomass
385 carriers (**Table S5**). The biomass carriers in those trickling filter reactors were exposed to
386 permanent shear forces, and shear forces changed microbial diversity and composition of the
387 developed biofilm (Rickard *et al.*, 2004) and promoted the proliferation of auto-aggregating
388 bacteria (Rochex *et al.*, 2008). In the nitrifying biofilm, AOB were preferentially localized on
389 the surface of the biofilm (*i.e.*, aerobic zone) whereas *Nitrospira* cells were easy to form cell
390 aggregates (Spieck *et al.*, 2006; Ushiki *et al.*, 2013) and heterologously distributed in the

391 biofilm and abundant in microaerobic zone (*i.e.*, inner part of biofilm) (Okabe *et al.*, 1999;
392 Schramm *et al.*, 2000). Such spatial distribution of AOB and NOB might occur in the sponge
393 carrier. Additionally, *Nitrospira* spp. in activated sludge tended to form physically stronger
394 cell aggregates than *Nitrosomonas oligotropha* (Larsen *et al.*, 2008), suggesting
395 *Nitrosomonas* in the NH₄⁺-feeding DHS reactor was likely to be more susceptible for the
396 detachment from biofilm. Occurrence of the washout of AOB and NOB from trickling filter
397 reactors has not been well investigated so far (Keuter *et al.*, 2011), which was examined in the
398 present study by determining the gene copy numbers of AOB *amoA*, *Nitrospira nxrB*, and
399 *Nitrospira* 16S rRNA genes (**Fig. 2**) in the NH₄⁺-enriched biomass and the reactor effluents.
400 The NMNS01 bin had one copy of *amoCAB* gene cluster as well as the closed relative
401 *Nitrosomonas* genomes (*Nitrosomonas* sp. Nm143, *Nitrosomonas* sp. UBA8640, and
402 *Nitrosomonas aestuarii*). As for *Nitrospira*, three *nxrAB* gene clusters and one 16S rRNA
403 gene were found in the *Nitrospira marina* genome. This ratio of the copy number of *nxrAB* to
404 16S rRNA gene (*i.e.*, 3) agreed with the ratio of the copy numbers of *Nitrospira nxrB* to
405 *Nitrospira* 16S rRNA gene found in the NH₄⁺-enriched biomass and the reactor effluents (*i.e.*,
406 3.0 to 3.4) (**Fig. 6**). Assuming one *amoA* gene copy per *Nitrosomonas* genome and three *nxrB*
407 and one 16S rRNA gene copy per *Nitrospira* genome, the abundance of *Nitrosomonas* was
408 1.5- and 1.7-folds greater (based on the normalized copy number of *nxrB* and 16S rRNA
409 gene, respectively) than *Nitrospira* in the effluents of NH₄⁺-feeding DHS reactor although

410 *Nitrospira* was 1.89- and 1.86-folds greater than *Nitrosomonas* in NH₄⁺-enriched biomass.
411 The greater abundance of *Nitrosomonas* in the effluents indicated that *Nitrosomonas* tended to
412 be washed out more frequently from the NH₄⁺-feeding DHS reactor. This washout of AOB
413 was another determinant for the dominance of *Nitrospira* over *Nitrosomonas* in the NH₄⁺-
414 feeding DHS reactors.

415 In summary, CO₂ assimilation efficiencies of *Nitrosomonas* and NO₂⁻-oxidizing
416 *Nitrospira* were determined, and the difference of the CO₂ assimilation efficiencies between
417 *Nitrosomonas* and *Nitrospira* was much smaller (0.077 μmol-CO₂/ μmol-NH₃ and 0.053–
418 0.054 μmol-CO₂/μmol-NO₂⁻, respectively) as compared with the difference of Δ*G_r*. Such
419 small difference in the CO₂ assimilation efficiencies was likely due to that large parts of free
420 energies during aerobic ammonia oxidation are dissipated and not conserved. The dissipation
421 of free energy (*i.e.*, efficiency of energy conservation) can not be expected from the value of
422 Δ*G_r*, and more bioenergetic studies, especially for *Nitrospira*, are required. *Nitrospira* use a
423 novel cytochrome *bd*-like heme-copper oxidase as a terminal oxidase and the nitrite oxidation
424 occurs in periplasmic spaces, which were not common with that of canonical NO₂⁻-oxidizing
425 *Nitrobacter*. It will be interesting to investigate the bioenergetic traits of *Nitrospira*. Apart
426 from the bioenergetics, washout of nitrifying population was another factor driving the
427 dominance of *Nitrospira* over *Nitrosomonas* in the DHS reactor; *i.e.*, *Nitrosomonas* was more
428 susceptible for washout than *Nitrospira*. Detachment and washout of particular nitrifiers has

429 been little explored, and the correlation between physicochemical parameters (*e.g.*, cell
430 surface hydrophobicity) remains to be explored in other studies.

431 **Experimental procedures**

432 **Operation of the DHS reactors fed with NH_4^+ or NO_2^-**

433 The 10-L DHS reactors (0.7 m in height and 0.17 m in width) were operated at 20°C in dark.
434 Details of the operated DHS reactors were previously described by the authors (Oshiki *et al.*,
435 2020). Briefly, the DHS reactors contained polyurethane-sponge media (183 pieces of cubic
436 sponge, 33 mm × 33 mm × 33 mm, set in a polypropylene tube, 32 mm diameter and 32 mm
437 long) as a biomass carrier. The sponge media had a 97% void ratio (*i.e.*, a percentage of the
438 volume of sponge pores), 256 m² m⁻³ of specific surface, and 0.63 mm of average pore size.
439 Artificial seawater media (pH 8.0, salinity 33‰) containing 0.297 g L⁻¹ NH₄Cl or 0.4 g L⁻¹
440 NaNO₂, 1 g L⁻¹ NaHCO₃, 34 g L⁻¹ artificial seawater powder (Marin Art, Tomita
441 Pharmaceutical, Naruto, Japan) was supplied to the top of the DHS reactor at the flow rate of
442 9.62 L d⁻¹ corresponding to 0.39 d of hydraulic retention time (HRT) and 200 mg-N L-sponge
443 media⁻¹ d⁻¹ of total ammonia nitrogen loading rate. The sponge media was exposed to the
444 atmosphere, and oxygen naturally dissolved into the media and aerobic condition was
445 maintained without external aeration. The filtrates were collected in a settling tank (volume;
446 0.9 L) located at the bottom of the DHS reactor, which were recirculated using a magnetic
447 pump at the flow rate of 4 L min⁻¹. The NH₄⁺- or NO₂⁻-feeding DHS reactors has been
448 operated for more than 1 y stably, and typical concentrations of NH₄⁺, NO₂⁻, NO₃⁻ and pH

449 values in the NH_4^+ -feeding DHS reactors were 7.9 μM , 9.3 μM , 4.9 mM, and pH 7.3,
450 respectively.

451 Biomass retained in the sponge media was collected by squeezing the sponge media
452 in the above artificial seawater media without NH_4Cl and NaNO_2^- , and subjected to the
453 following DNA extraction and batch incubations. As for the reactor effluents, two liters of the
454 effluents was daily collected from the NH_4^+ -feeding reactor for 4 d, and the collected
455 effluents were filtered on a 0.2 μm -pore-size PVDF membrane (Advantec, Tokyo, Japan)
456 separately. The filtered membranes were subjected to the DNA extraction.

457 **DNA extraction and determination of DNA concentration**

458 Genomic DNA was extracted from the biomass and the filtered membranes using a DNeasy
459 PowerSoil kit (Qiagen K.K., Tokyo, Japan) as following the instruction manual supplied by
460 manufactures. The concentrations of the extracted DNAs were determined using the Qubit
461 dsDNA BR assay kit and a Qubit 3.0 fluorospectrometer (Thermo Fisher Scientific, Waltham,
462 MD, USA).

463 **Metagenomic analysis**

464 A shotgun sequence library was prepared using a MGIEasy FS DNA Library Prep Set,
465 MGIEasy Circularization kit, and DNBSEQ-G400RS High-throughput sequencing set (MGI
466 Tech Japan, Tokyo, Japan). The 200-bp paired-end sequencing was performed using a
467 DNBSEQ-G400 sequencer. The paired-end sequence reads were trimmed using Trimmomatic

468 0.39 (SLIDINGWINDOW:6:30 MINLEN:100) (Bolger *et al.*, 2014). Digital normalization of
469 trimmed sequences was performed using bbnorm.sh of BBtools version 38.18 (target=100,
470 min=5) (<https://jgi.doe.gov/data-and-tools/bbtools/>). Assembled contigs were obtained from
471 NH₄⁺- and NO₂⁻-enriched biomass samples (co-assembly) by Megahit v1.2.9 (--k-min 27 --k-
472 max 141 --k-step 12) (Li *et al.*, 2015). Reads of each sample were mapped to assembled
473 contigs using bbmap.sh of BBtools. Obtained contigs of short length (< 2,500 bp) were
474 removed before binning. The multiple software of MaxBin2 version 2.15 (-markerset 40) (Wu
475 *et al.*, 2016), Metabat2 version 2.2.7 (Kang *et al.*, 2019), MyCC (MyCC_2017.ova) (Lin and
476 Liao, 2016) were used for binning from the contigs. To refine the obtained bins, we used
477 Binning_refiner version 1.4.0 with default parameters (Song and Thomas, 2017). The quality
478 of refined bins was checked using CheckM version 1.0.7 (Parks *et al.*, 2014). The relative
479 abundance of obtained bins was calculated using CoverM version 0.6.1
480 (<https://github.com/wwood/CoverM#installation>). Phylogenetic position of each bin is
481 estimated using GTDBtk v1.3.0 (release95) (Chaumeil *et al.*, 2019). Average nucleotide
482 identity (ANI) values of the obtained *Nitrospira* bins and the genomes in *Nitrospira*
483 sublineage IV were calculated using pyani version 0.2.11 (-m ANIb) (Pritchard *et al.*, 2016).
484 Gene prediction and annotation was performed via the D-FAST pipeline (Tanizawa *et al.*,
485 2018), and the MetaGeneAnnotator and Glimmer version 2.10, tRNAScan-SE version 1.23,

486 and blastn software applications were used for prediction of gene-coding sequences (CDSs),
487 tRNA, and rRNA, respectively.

488 Genomic tree was constructed using 120 concatenated phylogenetic marker genes of
489 obtained bins and representatives of genus *Nitrospira* or *Nitrosomonas* in the release95 of
490 GTDBtk version 1.3.0. For the multiple sequence alignment of *Nitrospira* spp., we included
491 the genomes of *Nitrospira marina* Nb-295T, *Nitrospirales* bacterium isolate MH-Pat-
492 all_autometa_1-10 (WLXC01000001), MAG-Baikal-G1, MAG-Baikal-deep-G158, MAG-
493 Baikal-deep-G159, MAG-ZH-13may13-77, MAG-cas150m-170, MAG-cas50m-175 with
494 *Nitrospira* genomes in the release95 of GTDBtk. Conserved marker genes were identified
495 using “gtdbtk identify” with default parameters and aligned to reference genomes using
496 “gtdbtk align” with taxonomic filters for phylogenies of *Nitrospira* (--taxa_fileter
497 c__Nitrospira), *Nitrosomonas* (--taxa_fileter f__Nitrosomonadaceae) or all metagenomic bins
498 (--taxa_fileter
499 f__UBA8639,g__Nitrosomonas,f__UBA11606,o__ARS69,f__Saprospiraceae,g__SZUA-
500 3,g__GCA-2699125,g__CR02bin9,f__Nitrospiraceae,f__SM1A02,g__UBA1845,f__B15-
501 G4,g__Hyphobacterium,g__Marinicaulis,g__UBA5701,g__Minwuia,f__Methyloligellaceae,f
502 __Rhodomicrobiaceae,g__UBA9145,o__UBA10353,g__UBA7359,o__Xanthomonadales).

503 Phylogenetic tree was constructed using IQ-TREE version 2.0.3 (-B 1000) with automatically
504 optimized substitution models (*Nitrospira*: LG+F+R7 and *Nitrosomonas*: JTT+F+R5) (Minh

505 *et al.*, 2020) and with the *Nitrospira lacus* (GCF_000355765.4) and *Thermodesulfovibrio*
506 *yellowstonii* (GCF_000020985.1) genomes for the *Nitrospira* and *Nitrosomonas* trees (**Fig. 2**
507 and **3**, respectively) as an outgroup.

508 **qPCR assay**

509 The qPCR assay was conducted using a ABI7500 fast Real-Time PCR System (Thermo
510 Fisher Scientific). The reaction mixture (10 μ L) contained KAPA SYBR FAST qPCR master
511 mix (Nippon Genetics, Tokyo, Japan) (5 μ L), 0.2 μ M each forward and reverse primer, 1 \times
512 ROX low dye, and 1 ng of the extracted DNA. Oligonucleotide primers used for PCR
513 amplification were 1) 515f and 806r for prokaryotic 16S rRNA gene, 2) amoA1F and
514 amoA2Rv1 for AOB *amoA* (Rotthauwe *et al.*, 1997, this study), 3) nxrB169f and nxrB638r
515 (Pester *et al.*, 2014) for *Nitrospira nxrB*, and 4) Nspra675F and Nspra746R (Graham *et al.*,
516 2007) for *Nitrospira* 16S rRNA gene, and the nucleotide sequences were shown in **Table S6**.
517 The original amoA2R primer (Rotthauwe *et al.*, 1997) had some mismatch bases against the
518 *amoA* sequence found in the NMNS01 bin, and the amoA2Rv1 was designed in the present
519 study by adding degenerate bases into the amoA2R primer. As for the amoA1F, nxrB169f,
520 nxrB638r, Nspra675F, and Nspra746R primers, there was no mismatch base between the
521 oligonucleotide primer and target gene found in the above metagenomic analysis. The cycling
522 conditions were the following: 95°C for 3 min; 40 cycles at 95°C for 3 s and 60°C for 20 s;
523 and, finally, 65°C to 95°C in 0.5°C increments for the melting curve analysis. Standard curves

524 (10^1 to 10^6 copies/ μL) were prepared using a dilution series of plasmid DNAs containing PCR
525 products of the above target. Partial sequences of *Escherichia coli* JM109 (SMOBIO
526 technology, Hsinchu, Taiwan) 16S rRNA gene, *Nitrosomonas europaea* (NBRC14298)
527 *amoA*, *Nitrospira inopinata* (JCM31988) 16S rRNA gene were amplified using the above
528 oligonucleotide primer set. *Nitrospira nxrB* was amplified using the genomic DNA extracted
529 from NO_2^- -enriched biomass. The obtained PCR products were cloned into pUC118 vector
530 using mighty cloning reagent (TakaraBio, Shiga, Japan), and transformed into *E. coli* DH5 α
531 cells (SMBIO technology) by heat shock. Plasmids were extracted from the transformants
532 using FastGene Plasmid mini kit (Nippon Genetics), nucleotide sequences of the cloned PCR
533 products were ascertained by performing the Sanger sequencing, and the concentrations of the
534 extracted plasmids were determined fluorometrically as previously described above.

535 Activity tests

536 Assimilation of $^{14}\text{CO}_2$ into the biomass during aerobic ammonia and nitrite oxidation
537 was examined as previously described (Oshiki *et al.*, 2011). Briefly, the 2.5 mL of biomass
538 suspension (24 and 0.66 μg -protein mL^{-1} of biomass concentrations for NH_4^+ - and NO_2^- -
539 enriched biomass, respectively) containing 0.5 mM NH_4^+ or NO_2^- was incubated at 20°C in
540 10-mL glass vials with shaking at 60 rpm. The ^{14}C -labeled sodium bicarbonate (Moravek Inc.,
541 Brea, CA, USA) was added at a concentration of 10 $\mu\text{Ci vial}^{-1}$, and the vials were sealed with
542 butyl rubber plug and aluminum seal. After 18 h of incubation, the biomass was fixed with

543 4% paraformaldehyde, washed three times with PBS, and mixed with scintillation cocktail
544 Clear-sol I (Nacalai, Tokyo, Japan). Radioactivity was determined with an ALOKA LSC-
545 6100 liquid scintillation counter. Additional cold incubation with ^{12}C -labeled sodium
546 bicarbonate instead of ^{14}C -labeled one was performed in parallel to determine ammonia and
547 nitrite oxidation rates. Abiotic incorporation of ^{14}C -labeled sodium bicarbonate was examined
548 by performing the above incubation with the biomass pasteurized at 70°C for 15 min. For the
549 incubation using H_2 as a substrate instead of NH_4^+ and NO_2^- , biomass suspension without
550 NH_4^+ and NO_2^- were dispensed into the closed vials, and pure H_2 gas (GL Science, Tokyo,
551 Japan) was injected into headspace at the final concentration of 2% (v/v) using a gas tight
552 syringe.

553 Chemical analysis

554 NH_4^+ concentration was determined fluorometrically using the *o*-phthalaldehyde (OPA)
555 method (Taylor *et al.*, 1974). Liquid samples were mixed with 3.8 mM *o*-phthalaldehyde, and
556 fluorescence intensity was determined at 355 nm of excitation and 460 nm of emission.

557 NO_2^- concentration was determined colorimetrically using the
558 naphthylethylenediamine method (Rice *et al.*, 2012). Liquid samples were mixed with a
559 naphthylethylenediamine-sulfanilamide solution, and absorbance was measured at 540 nm.

560 NO_3^- concentration was determined colorimetrically using the brucine sulfate
561 method (Jenkins and Medsker, 1964). Liquid samples were mixed with 80% (vol/vol) sulfuric

562 acid and brucine sulfanilic acid solution, and heated at 100°C for 20 min. The absorbance was
563 measured at 410 nm.

564 Protein concentration was determined by the Lowry method using a DC protein
565 assay kit (Bio-Rad, Hercules, CA, USA). Bovine serum albumin was used as a protein
566 standard.

567 H₂ concentration was determined by gas chromatography as described elsewhere.
568 The 100 µL of gas sample was injected into a gas chromatograph GC-2014 equipped with a
569 thermal conductivity detector and a 2-m stainless column packed with a Molecular Sieve-5A.

570 **Fluorescence in-situ hybridization and microscopy**

571 Fixation of biomass (4% paraformaldehyde) and *in situ* hybridization of oligonucleotide
572 probes were performed as previously described (Kindaichi *et al.*, 2004). The fixed biomass
573 was sonicated at 3 watt for 4 minutes, and hybridized with the oligonucleotide probe
574 Ntspa712 (Daims *et al.*, 2001) for *Nitrospira* or Nsm156 (Mobarry *et al.*, 1996) for
575 *Nitrosomonas*, respectively.

576 **Calculation of CO₂ assimilation and free-energy efficiencies.**

577 CO₂ assimilation efficiencies of NH₄⁺- and NO₂⁻-enriched biomass during aerobic ammonia
578 and nitrite oxidation (unit; µmol-CO₂/µmol-NH₃ or µmol-NO₂⁻) were calculated by dividing
579 the molar amounts of ¹⁴CO₂ fixed during the incubation with those of the consumed NH₄⁺ and
580 NO₂⁻. The fixed CO₂ includes the carbon fixed into cellular materials and also that fixed as

581 extracellular polymeric substances (EPS) (Okabe *et al.*, 2005). On the other hand, the fixed
582 CO₂ does not include the carbon released as dissolved organic carbon (DOC); *i.e.*, *ca.* 6-8%
583 and 12% of fixed CO₂ were released as DOC in the culture of marine AOB (*Nitrosomonas*
584 *marina* C-25 and *Nitrosomonas* sp. C-15) and marine NOB (*Nitrospira marina* Nb-295),
585 respectively (Bayer *et al.*, 2022).

586 The values of free-energy efficiency were calculated using the following equation
587 (Glover, 1985).

$$588 \quad \text{free - energy efficiency (\%)} = \text{growth yield} \times \frac{495}{\Delta G_r} \times 100$$

589 Where, 495; the free energy (kJ/mol) required for converting CO₂ into CH₂O, and ΔG_r ; the
590 free energy (kJ/mol) obtained from aerobic ammonia or nitrite oxidation during the batch
591 incubations (*i.e.*, 283.3 kJ /mol-NH₃ and 83.5 kJ /mol-NO₂⁻, respectively) (**Supplementary**
592 **text 1**). For AOB and NOB examined in the previous studies, the 286.7 kJ /mol-NH₃ for
593 AOB, and 73.8 kJ /mol-NO₂⁻ for NOB were used. Those values corresponded to ΔG_r^m that
594 were the Gibbs free energy change at when the concentrations of all reactants were 1 mM at
595 pH 7 and 25°C (**Supplementary text 1**).

596 **Accession numbers**

597 Raw metagenomic sequence data and the assembled and annotated 30 bins obtained in the
598 present study are available in the DDBJ nucleotide sequence database under the accession
599 number DRA013035 and those in **Table S7**, respectively.

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610 **Conflict of interest**

611 The authors declare no conflicts of interest associated with this manuscript.

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832 **Table 1. Summary of metagenomic bins.** *Nitrospira*, *Nitrosomonas*, and *Nitrospina* bins

833 were designated with the label of “NPIRA”, “NMNS”, and “NPINA”, respectively.

834 Phylogeny of the other DHS20C bins were shown in **Fig. S1**.

Bin(s)	Relative abundance (%)		Size (Mb)	Contigs	CDSs	Completeness	Contamination	Strain heterogeneity
	NH ₄ ⁺ - enriched	NO ₂ ⁻ - enriched						
NPIRA01	1.44	39.7	4.7	203	4,095	90%	3%	50
NPIRA02	35.0	21.2	4.7	92	4,275	97%	3%	33
NPIRA03	1.08	1.50	4.9	218	4,203	94%	4%	20
NPIRA04	14.1	8.88	4.8	633	3,683	81%	4%	30
NPIRA05	0.59	0.73	3.1	595	2,336	59%	0%	0
NPIRA06	1.52	0.21	4.0	303	3,436	85%	5%	57
NMNS01	1.72	0.58	3.7	199	3,093	98%	1%	0
NMNS02	5.86	0.04	3.6	117	3,117	99%	2%	0
NPINA01	0.13	2.10	3.6	14	3,355	97%	3%	0
DHS20C01	1.01	0.73	4.4	119	3,859	99%	3%	8
DHS20C02	0.00	3.43	2.1	12	2,066	96%	1%	0
DHS20C03	1.47	0.51	4.6	234	4,050	100%	2%	82
DHS20C04	1.63	0.00	3.4	133	3,254	95%	0%	0

Bin(s)	Relative abundance (%)		Size (Mb)	Contigs	CDSs	Completeness	Contamination	Strain heterogeneity
	NH ₄ ⁺ - enriched	NO ₂ ⁻ - enriched						
DHS20C05	0.43	0.00	3.0	386	2,588	84%	1%	17
DHS20C06	0.46	0.10	2.4	271	2,137	88%	0%	0
DHS20C07	2.79	0.07	3.6	27	3,178	100%	1%	0
DHS20C08	2.41	0.75	2.9	6	2,510	98%	1%	0
DHS20C09	0.46	0.07	2.8	420	2,257	84%	8%	3
DHS20C10	1.04	0.03	1.7	65	1,458	89%	1%	67
DHS20C11	0.00	0.81	4.7	210	3,873	96%	1%	0
DHS20C12	0.55	0.03	3.6	282	3,076	91%	11%	39
DHS20C13	1.01	0.14	3.4	173	3,114	97%	7%	9
DHS20C14	0.61	0.27	2.9	273	2,298	91%	3%	33
DHS20C15	1.41	0.03	4.5	11	3,500	96%	1%	0
DHS20C16	0.50	0.02	5.3	478	3,780	96%	9%	0
DHS20C17	0.46	0.19	5.2	462	3,634	97%	3%	0
DHS20C18	0.83	0.05	8.0	161	5,634	100%	1%	0
DHS20C19	0.44	0.17	3.7	589	3,085	80%	3%	0
DHS20C20	0.01	0.68	4.9	1044	3,440	73%	15%	70

Bin(s)	Relative abundance (%)		Size (Mb)	Contigs	CDSs	Completeness	Contamination	Strain heterogeneity
	NH ₄ ⁺ - enriched	NO ₂ ⁻ - enriched						
DHS20C21	0.01	0.69	3.6	653	2,475	77%	4%	0

835

836 **Table 2 CO₂ assimilation and free-energy efficiencies of NH₄⁺- and NO₂⁻-enriched**
837 **biomass, AOB, and NOB.** CO₂ assimilation efficiencies examining ¹⁴CO₂ fixation into
838 biomass during NH₃ or NO₂⁻ oxidation are summarized here, and the values are available as
839 (mean ± standard deviations). NA; not available because the NO₂⁻-enriched biomass did not
840 show the activity of aerobic ammonia oxidation. *; the value was calculated by subtracting the
841 CO₂ assimilation efficiency of aerobic ammonia oxidation with that of nitrite oxidation. The
842 CO₂ assimilation efficiencies in this table does not include a fraction of the carbon released as
843 dissolved organic carbon (DOC). The reference data of Bayer *et al.* (2022) are a personal gift
844 from Dr. Barbara Bayer.

Biomass/microorganisms	Reaction	μmol-CO ₂ /μmol-NH ₃ or NO ₂ ⁻ (Free-energy efficiency)	Reference
NH ₄ ⁺ -enriched biomass	NH ₃ → NO ₃ ⁻	0.13 ± 0.019	this study
	NH ₃ → NO ₂ ⁻	0.077* (13%)	this study
	NO ₂ ⁻ → NO ₃ ⁻	0.053 ± 0.013 (31%)	this study
NO ₂ ⁻ -enriched biomass	NH ₃ → NO ₂ ⁻	NA	this study
	NO ₂ ⁻ → NO ₃ ⁻	0.054 ± 0.019 (32%)	this study
AOB			
<i>Nitrosomonas marina</i>	NH ₃ → NO ₂ ⁻	0.04 – 0.07 (7 – 12%)	Glover (1985)
<i>Nitrosococcus oceanus</i>	NH ₃ → NO ₂ ⁻	0.024 – 0.062 (4 – 11%)	Glover <i>et al.</i> (1985)
<i>Nitrosomonas marina</i> C-25	NH ₃ → NO ₂ ⁻	0.043 ± 0.012 (7%)	Bayer <i>et al.</i> (2022)

Biomass/microorganisms	Reaction	$\mu\text{mol-CO}_2/\mu\text{mol-NH}_3$ or NO_2^- (Free-energy efficiency)	Reference
<i>Nitrosomonas sp. C-15</i>	$\text{NH}_3 \rightarrow \text{NO}_2^-$	0.044 ± 0.007 (8%)	Bayer <i>et al.</i> (2022)
NOB			
<i>Nitrococcus mobilis</i>	$\text{NO}_2^- \rightarrow \text{NO}_3^-$	0.014 - 0.031 (9 – 21%)	Glover (1985)
<i>Nitrococcus mobilis Nb-231</i>	$\text{NO}_2^- \rightarrow \text{NO}_3^-$	0.016 ± 0.002 (11%)	Bayer <i>et al.</i> , 2022
<i>Nitrobacter agilis</i>	$\text{NO}_2^- \rightarrow \text{NO}_3^-$	0.009 (6%)	Tsai and Tuovinen (1986)
<i>Nitrospira marina Nb-295</i>	$\text{NO}_2^- \rightarrow \text{NO}_3^-$	0.032 ± 0.005 (22%)	Bayer <i>et al.</i> , 2022
<i>Nitrospina sp. Nb-3</i>	$\text{NO}_2^- \rightarrow \text{NO}_3^-$	0.035 ± 0.005 (23%)	Bayer <i>et al.</i> , 2022
<i>Nitrospina gracilis Nb-211</i>	$\text{NO}_2^- \rightarrow \text{NO}_3^-$	0.029 ± 0.002 (19%)	Bayer <i>et al.</i> , 2022

845

846 **Figure legends**

847 **Fig. 1. Fluorescence *in-situ* hybridization analysis of NH₄⁺-enriched biomass.** The
848 biomass was fixed with 4% paraformaldehyde, hybridized with the oligonucleotide probe
849 Ntspa712 (labeled with Cy3) for *Nitrospira* (panel **a**) or Nsm156 (TRITC) for *Nitrosomonas*
850 (panel **b**), and stained with DAPI. The cells showing red and cyan color represent *Nitrospira*
851 or *Nitrosomonas* population (red) and total cells (cyan), respectively. Bar = 20 μm.

852 **Fig. 2. Genome tree showing the phylogeny of *Nitrospira* bins.** A phylogenetic clade of
853 *Nitrospira* sublineage IV was shown with a bracket, and the phylogenetic position of the
854 obtained *Nitrospira* bins affiliated into *Nitrospira marina* and *Nitrospira salsa* clades were
855 shown with red color. The scale bar represents 10% sequence divergence.

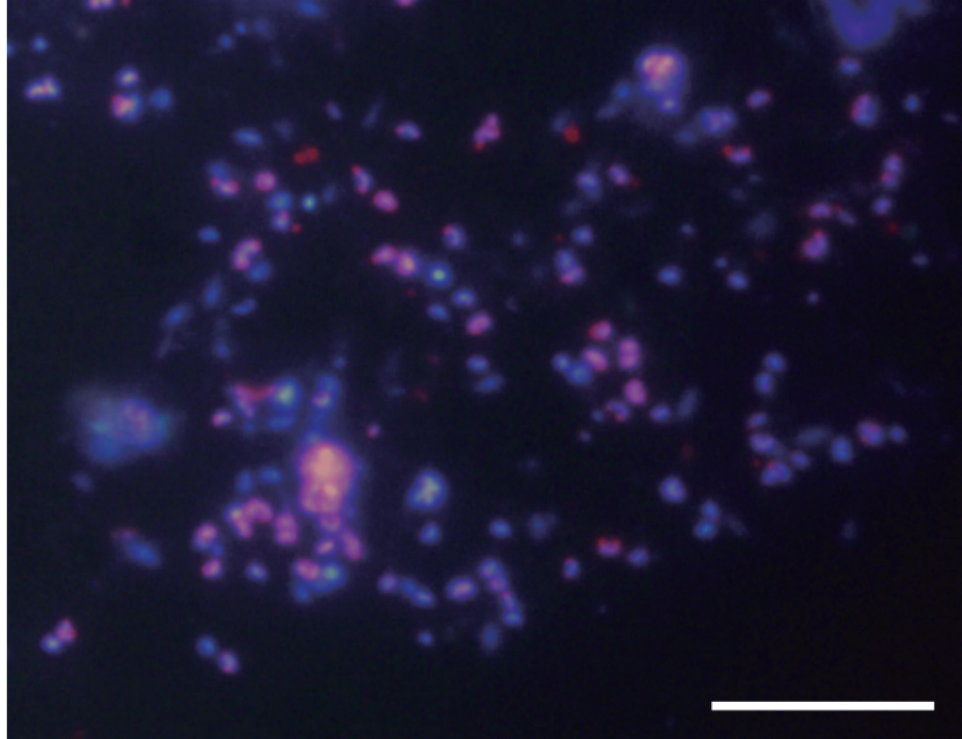
856 **Fig. 3. Genome tree showing the phylogeny of *Nitrosomonas* bins.** The phylogenetic
857 position of the obtained *Nitrosomonas* bins were shown with red color. The scale bar
858 represents 5% sequence divergence.

859 **Fig. 4. Nitrification activities of NH₄⁺- and NO₂⁻-enriched biomass (left and right panels,**
860 **respectively).** The biomass was aerobically incubated with the addition of NH₄⁺ or NO₂⁻, and
861 the concentrations of NH₄⁺, NO₂⁻, and NO₃⁻ were determined. The incubation was performed
862 in triplicates, and the symbol and error bars represent the mean value and the range of
863 standard deviation, respectively.

864 **Fig. 5. Aerobic H₂ oxidation by NH₄⁺- and NO₂⁻-enriched biomass.** The biomass
865 suspension (2.5 mL) was incubated in closed 10-mL vials with the addition of pure H₂ gas
866 into the head space, and the concentrations of H₂ and O₂ in the head space were monitored.
867 Both the biomass consumed H₂, while the consumption required more than 4 d of lag phase.
868 Error bars represent the range of standard deviation derived from triplicate incubations.

869 **Fig. 6. Abundance of AOB and *Nitrospira* in the NH₄⁺-enriched biomass and the reactor**
870 **effluents discharged from the NH₄⁺-feeding DHS reactor.** Copy numbers of bacterial 16S
871 rRNA gene, AOB *amoA*, *Nitrospira nxrB*, and *Nitrospira* 16S rRNA gene were determined
872 by quantitative PCR assay. Genomic DNA extraction was performed with >3 biomass
873 samples, and error bar represent the standard deviation of the copy numbers determined from
874 each DNA extracts.

a)



b)

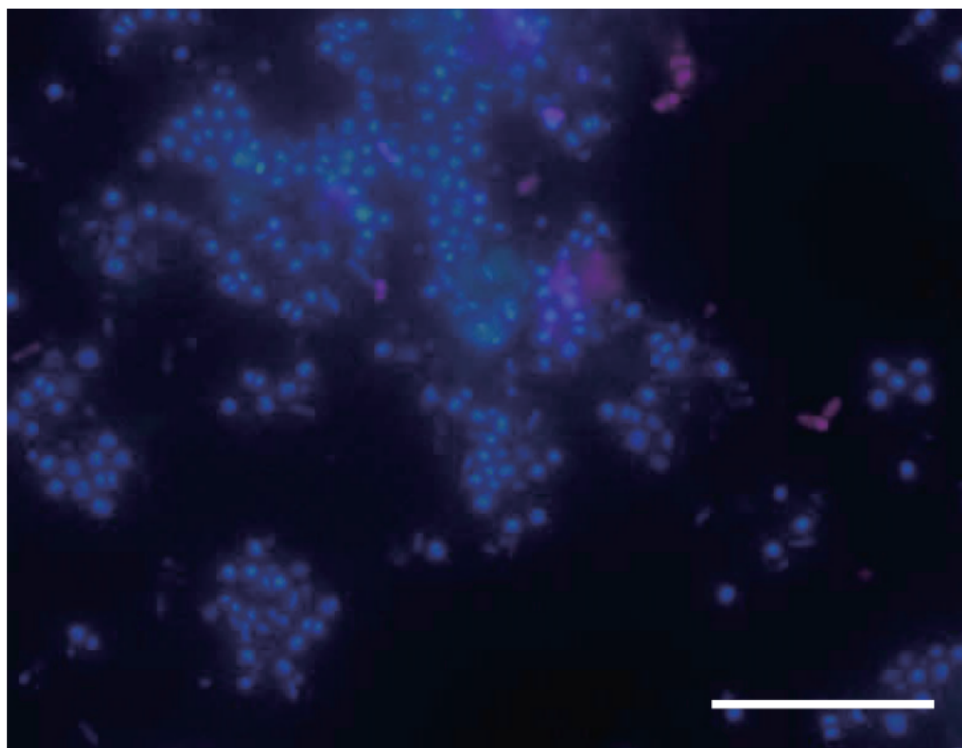


Fig. 1 (Oshiki et al.)

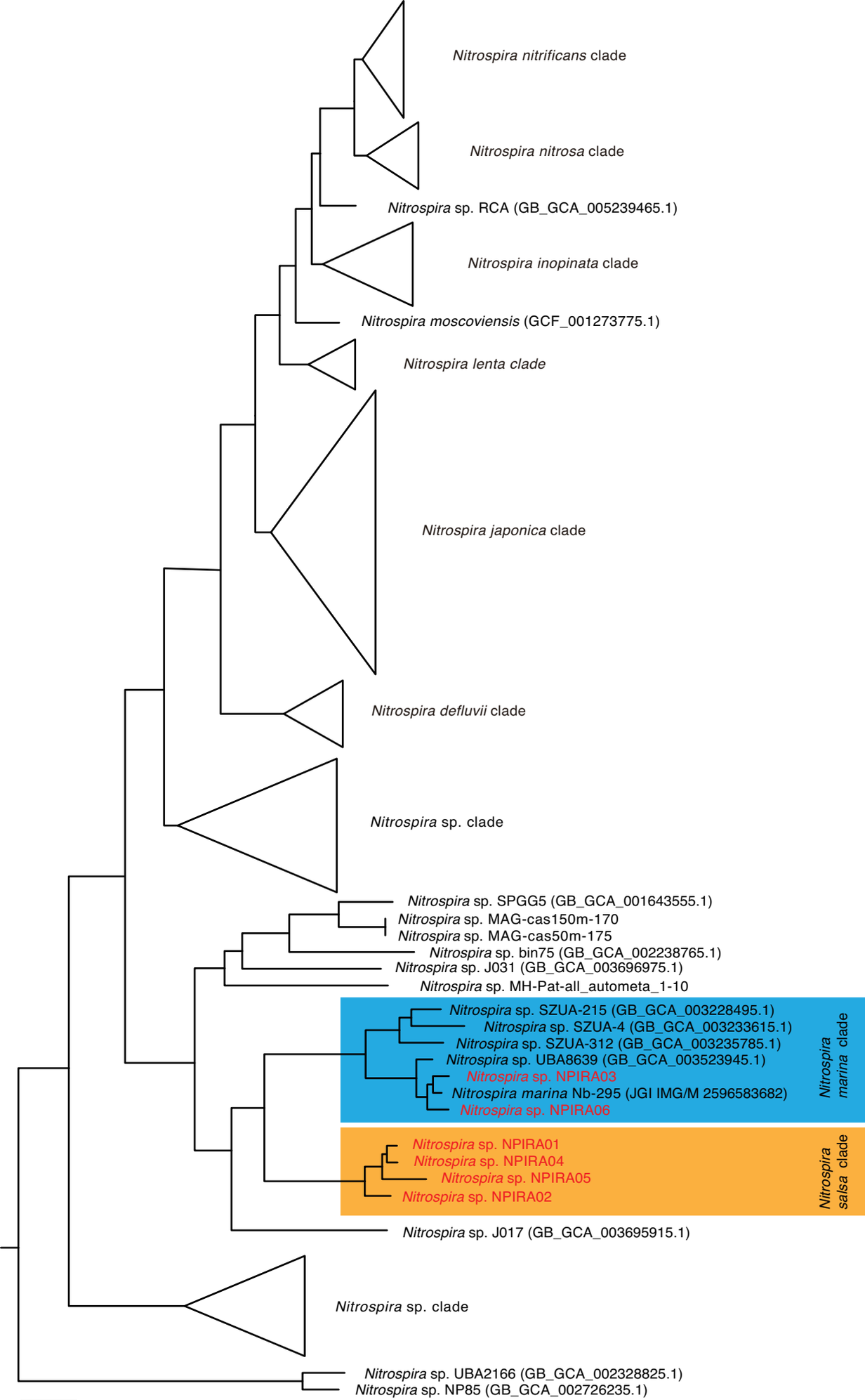


Fig. 2 (Oshiki et al.)

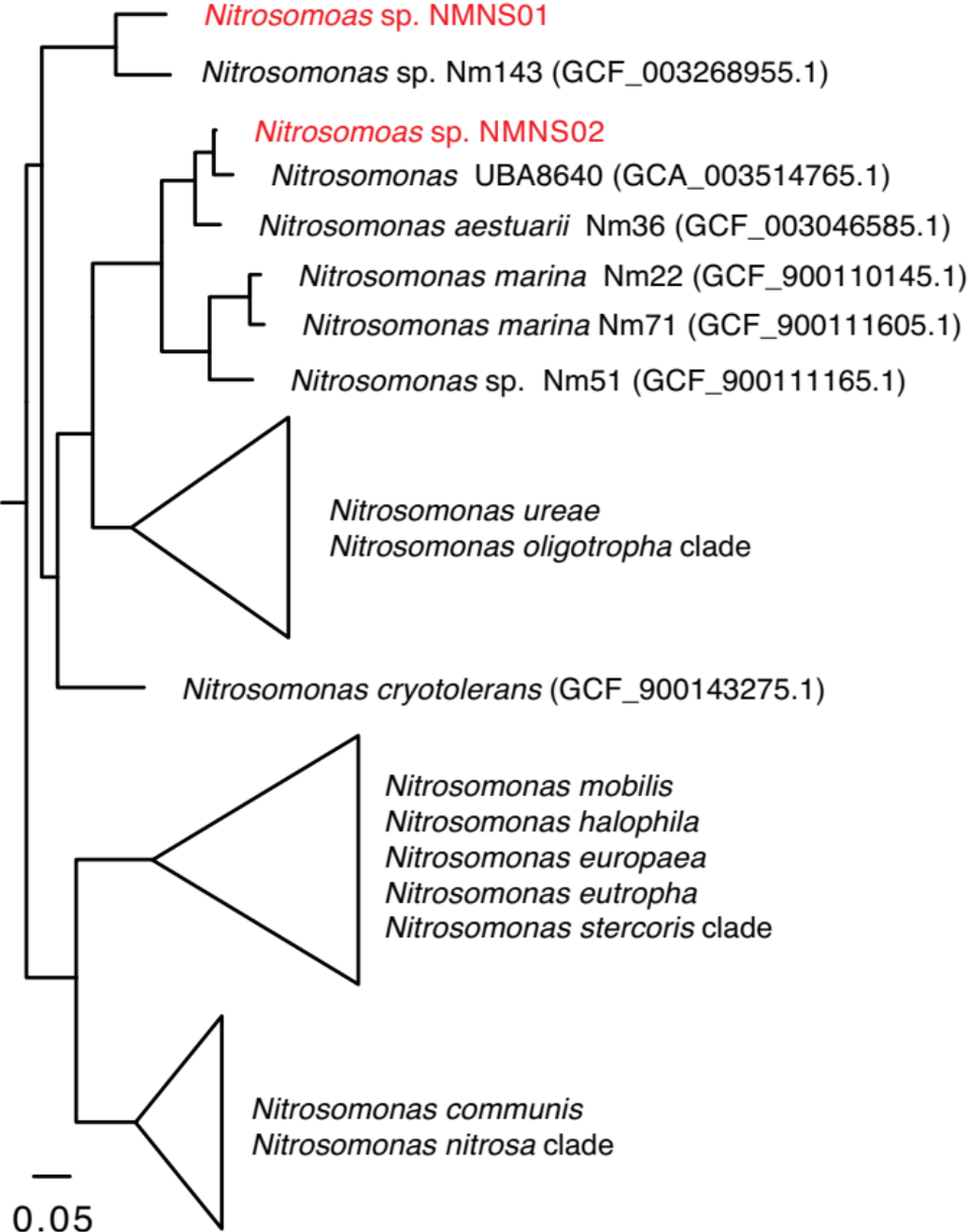


Fig. 3 (Oshiki et al.)

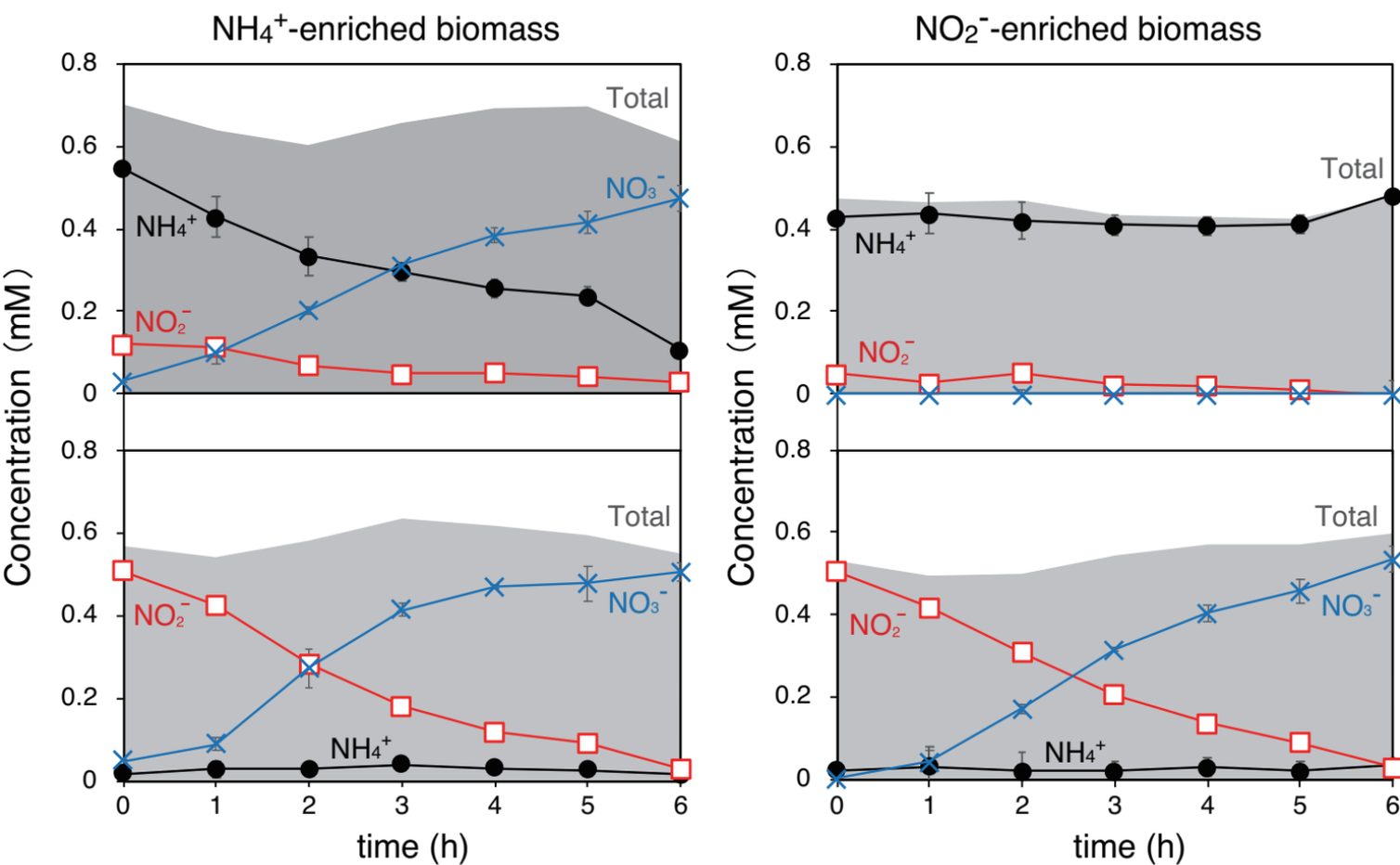
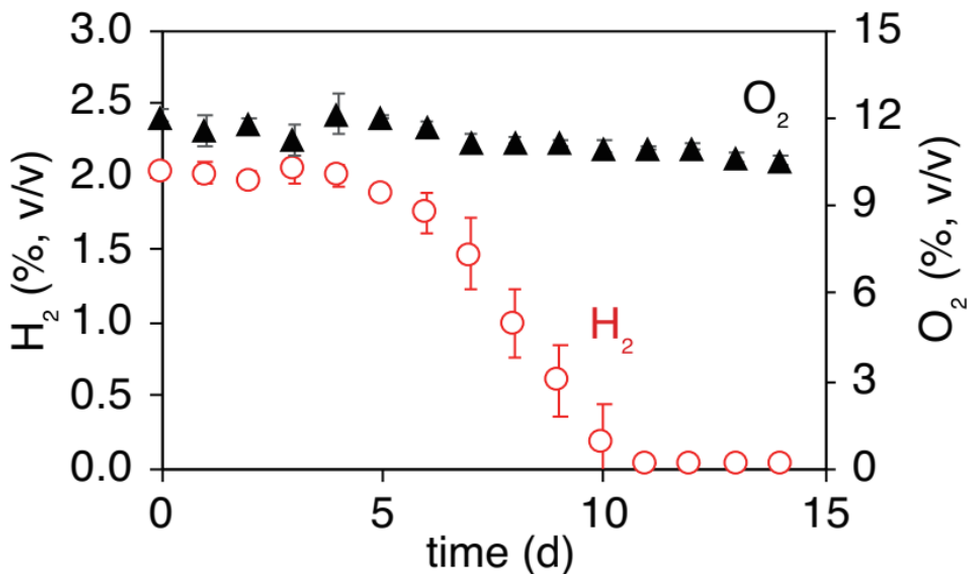


Fig. 4 (Oshiki et al.)

a) NH_4^+ -enriched biomass



b) NO_2^- -enriched biomass

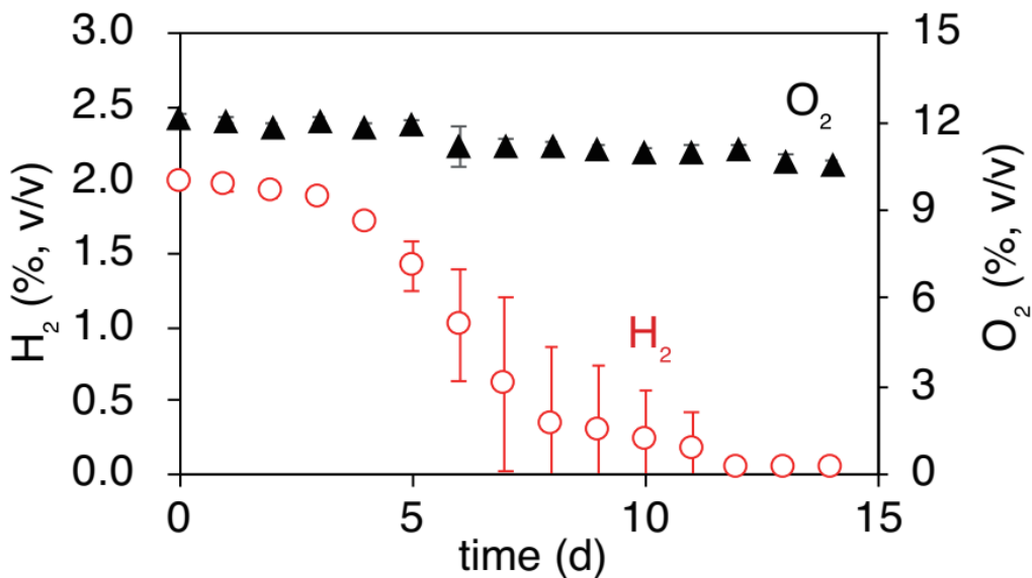


Fig. 5 (Oshiki et al.)

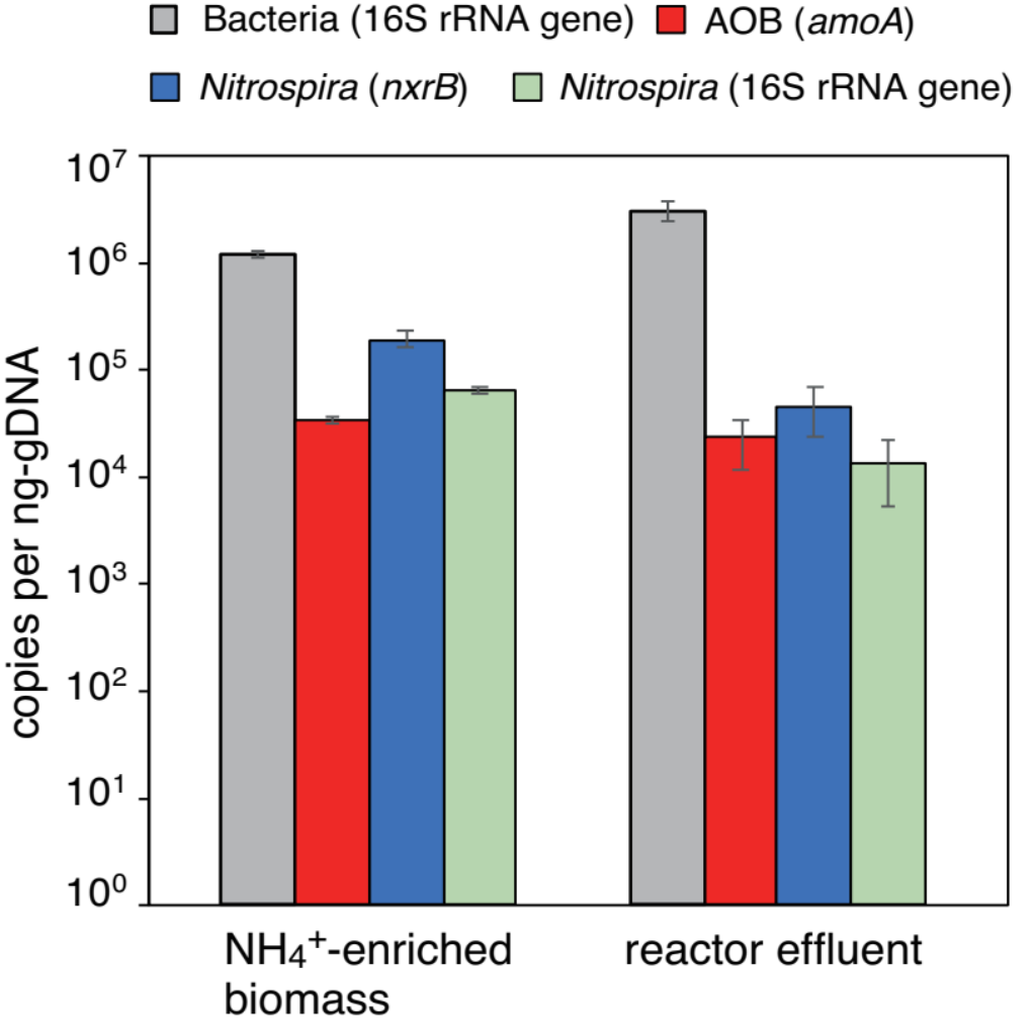


Fig. 6 (Oshiki et al.)